

# Package ‘hpoPlot’

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**Type** Package

**Title** Functions for Plotting HPO Terms

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**Description** Collection of functions for manipulating sets of HPO terms and plotting them with a various options.

**License** GPL (>= 2)

**Depends** R (>= 3.0.0)

**Imports** methods, Rgraphviz, functional, magrittr

**Suggests** knitr

**VignetteBuilder** knitr

**RoxygenNote** 5.0.1

**NeedsCompilation** no

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---

apply.term.filters      *Apply a list of term filters to a given plotting context*

---

**Description**

Apply a list of term filters to a given plotting context

**Usage**

```
apply.term.filters(hpo.terms, plotting.context, term.filters,
  starting.terms = NULL)
```

**Arguments**

hpo.terms      R-Object representation of HPO

plotting.context      List object with hpo.phenotypes slot for list of character vectors of terms

term.filters      List of term filtering functions

starting.terms      Character vector of HPO term codes to filter. Defaults to all terms in the 'hpo.phenotypes' element of plotting.context, if it is present

**Value**

Character vector of terms

**Examples**

```
data(hpo.terms)
apply.term.filters(hpo.terms=hpo.terms, plotting.context=list(
  hpo.phenotypes=list(Case1="HP:0001873")), term.filters=list(remove.links))
```

---

calibrate.sizes      *Function to scale sizes of terms between two given limits*

---

**Description**

Function to scale sizes of terms between two given limits

**Usage**

```
calibrate.sizes(x, high, low)
```

**Arguments**

x              Numeric vector of term relative sizes named by term

high            Numeric vector of largest size

low             Numeric vector of smallest size

**Value**

Numeric vector

**Examples**

```
calibrate.sizes(c("HP:0000001"=10, "HP:0000006"=5), high=3, low=1)
```

---

clean.terms	<i>Remove redundant/implied terms</i>
-------------	---------------------------------------

---

**Description**

Remove redundant/implied terms

**Usage**

```
clean.terms(hpo.terms, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms

**Value**

Character vector of HPO terms

**Examples**

```
data(hpo.terms)
clean.terms(hpo.terms, c("HP:0001873", "HP:0001872"))
```

---

exclude.branch	<i>Exclude terms descending from particular term from a character vector of terms</i>
----------------	---

---

**Description**

Exclude terms descending from particular term from a character vector of terms

**Usage**

```
exclude.branch(hpo.terms, branch.root, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
branch.root	HPO term whose descendants should be excluded
terms	Character vector of HPO terms

**Value**

Character vector of terms

---

get.ancestors	<i>Get set of all ancestors of set of terms</i>
---------------	---

---

**Description**

Get set of all ancestors of set of terms

**Usage**

```
get.ancestors(hpo.terms, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms

**Value**

Character vector of all HPO terms which are an ancestor of at least one term in terms, including the terms themselves

**See Also**

link{get.descendants}

**Examples**

```
data(hpo.terms)
get.ancestors(hpo.terms, c("HP:0001873", "HP:0011877"))
```

---

get.case.based.colours
------------------------

*Function to set colours of HPO nodes in plot to distinguish terms belonging to different sets of phenotypes*

---

**Description**

Function to set colours of HPO nodes in plot to distinguish terms belonging to different sets of phenotypes

**Usage**

```
get.case.based.colours(hpo.terms, terms, plotting.context)
```

**Arguments**

hpo.terms      R-Object representation of HPO  
 terms          Character vector of HPO terms  
 plotting.context  
                  List object with hpo.phenotypes slot for list of character vectors of terms

**Value**

Character vector of colours, named by term

---

get.case.based.labels    *Function to label HPO nodes in plot to indicate to which phenotypes each of the terms belong*

---

**Description**

Function to label HPO nodes in plot to indicate to which phenotypes each of the terms belong

**Usage**

```
get.case.based.labels(hpo.terms, terms, plotting.context)
```

**Arguments**

hpo.terms      R-Object representation of HPO  
 terms          Character vector of HPO terms  
 plotting.context  
                  List object with hpo.phenotypes slot for list of character vectors of terms

**Value**

Character vector of colours, named by term

---

get.case.term.matrix    *Get a matrix with columns of hpo terms and rows of patients,*

---

**Description**

Get a matrix with columns of hpo terms and rows of patients,

**Usage**

```
get.case.term.matrix(hpo.phenotypes, columns = NULL)
```

**Arguments**

- `hpo.phenotypes` List of character vectors of HPO terms. Result includes only terms which are explicitly present in the list items, so if you wish the result to include even terms which are implicitly present, lapply [get.ancestors](#) to the argument before passing it to this function
- `columns` Force result to have these exact columns, entering F for terms which aren't present

**Value**

Logical matrix - entry for a patient/hpo term = T if the patient has the term and F otherwise.

**Examples**

```
get.case.term.matrix(list(Patient1=c("HP:0001873")))
```

---

`get.code.node.labels` *Function to label HPO nodes in plot with just HPO code*

---

**Description**

Function to label HPO nodes in plot with just HPO code

**Usage**

```
get.code.node.labels(hpo.terms, terms, plotting.context)
```

**Arguments**

- `hpo.terms` R-Object representation of HPO
- `terms` Character vector of HPO terms
- `plotting.context` List object with `hpo.phenotypes` slot for list of character vectors of terms

**Value**

Character vector of labels, named by term

---

get.descendants      *Get set of all descendants of single term*

---

**Description**

Get set of all descendants of single term

**Usage**

```
get.descendants(hpo.terms, ancestor, remove.ancestor = FALSE)
```

**Arguments**

hpo.terms	R-Object representation of HPO
ancestor	Character vector of length 1 - the HPO code of the term whose descendants you wish to retrieve
remove.ancestor	Boolean indicating whether to remove the given ancestor or not

**Value**

Character vector of terms

**See Also**

link{get.ancestors}

**Examples**

```
data(hpo.terms)
get.descendants(hpo.terms, ancestor=c("HP:0001873"))
```

---

get.frequency.based.colours  
*Function to colour HPO nodes in plot with colours based on frequency with which terms appear in phenotypes*

---

**Description**

Function to colour HPO nodes in plot with colours based on frequency with which terms appear in phenotypes

**Usage**

```
get.frequency.based.colours(hpo.terms, terms, plotting.context,
  colour.func = NULL)
```

**Arguments**

<code>hpo.terms</code>	R-Object representation of HPO
<code>terms</code>	Character vector of HPO terms
<code>plotting.context</code>	List object with <code>hpo.phenotypes</code> slot for list of character vectors of terms
<code>colour.func</code>	Function capable of returning a set of colours, given the number of colours it needs to return

**Value**

Character vector of colours, named by term

---

`get.frequency.based.labels`

*Function to label HPO nodes in plot based on frequency of occurrence in phenotypes*

---

**Description**

Function to label HPO nodes in plot based on frequency of occurrence in phenotypes

**Usage**

`get.frequency.based.labels(hpo.terms, terms, plotting.context)`

**Arguments**

<code>hpo.terms</code>	R-Object representation of HPO
<code>terms</code>	Character vector of HPO terms
<code>plotting.context</code>	List object with <code>hpo.phenotypes</code> slot for list of character vectors of terms

**Value**

Character vector of labels, named by term

get.frequency.based.sizes

*Function to size HPO nodes in plot based on frequency of occurrence in phenotypes*

---

**Description**

Function to size HPO nodes in plot based on frequency of occurrence in phenotypes

**Usage**

```
get.frequency.based.sizes(hpo.terms, terms, plotting.context)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms

**Value**

Character vector of sizes, named by term

---

get.full.labels

*Function to label HPO nodes in plot with full labels*

---

**Description**

Function to label HPO nodes in plot with full labels

**Usage**

```
get.full.labels(hpo.terms, terms, plotting.context)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms

**Value**

Character vector of labels, named by term

---

get.hpo.graph	<i>Get HPO graph object</i>
---------------	-----------------------------

---

**Description**

Get HPO graph object

**Usage**

```
get.hpo.graph(hpo.terms, terms = apply.term.filters(hpo.terms = hpo.terms,
  plotting.context = plotting.context, term.filters = list()),
  plotting.context = NULL, colours = "white",
  labels = get.simple.node.labels, borders = "#FFFFFF00", sizes = 0.75,
  font.sizes = rep(30, length(terms)), shapes = rep("circle",
  length(terms)), nodeAttrs = NULL, arrowAttrs = list(color = "#000000"))
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms
colours	Function to set the colours of the HPO nodes in the graph based on the plotting context, or a character vector of colours
labels	Function to set the labels of the HPO nodes in the graph based on the plotting context, or a character vector of node labels
borders	Function to set the borders of the HPO nodes in the graph based on the plotting context, or a character vector of border colours
sizes	Function to set the sizes of the HPO nodes in the graph based on the plotting context, or a numeric vector of node sizes
font.sizes	Function to set the font sizes of the text to be placed in the HPO nodes in the graph based on the plotting context, or an integer vector of font sizes
shapes	Function to set the shapes of the HPO nodes in the graph based on the plotting context, or a character vector of shape names (defaults to 'circle')
nodeAttrs	Pass nodeAttrs directly to rgraphviz plotting function
arrowAttrs	List of properties to set for arrows (note, these properties will be used for all arrow)

**Value**

graphAM S4 object

**See Also**

[hpo.plot](#)

## Examples

```
data(hpo.terms)
phenotype.strings <- c(
  A="HP:0001382,HP:0004272,HP:0007917,HP:0004912,HP:0001596",
  B="HP:0001382,HP:0004272,HP:0002165,HP:0004800,HP:0004912",
  C="HP:0004800,HP:0001382,HP:0004912,HP:0007917,HP:0008743",
  D="HP:0001257,HP:0001382,HP:0007917,HP:0012623,HP:0002165",
  E="HP:0007917,HP:0004800,HP:0004272,HP:0001596,HP:0002165"
)

hpo.phenotypes <- term.set.list.from.character(phenotype.strings)

get.hpo.graph(
  hpo.terms=hpo.terms,
  plotting.context=list(hpo.phenotypes=hpo.phenotypes)
)
```

---

get.informative.node.labels

*Function to label HPO nodes in plot with node description and information content*

---

## Description

Function to label HPO nodes in plot with node description and information content

## Usage

```
get.informative.node.labels(hpo.terms, terms, plotting.context)
```

## Arguments

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms

## Value

Character vector of labels, named by term

---

get.mpo.to.hpo            *Get MPO to HPO R-Object*

---

**Description**

Get MPO to HPO R-Object

**Usage**

```
get.mpo.to.hpo(hpo.terms, cross.species.file)
```

**Arguments**

hpo.terms	R-Object representation of HPO
cross.species.file	cross species obo file, downloadable from <a href="http://compbio.charite.de/hudson/">http://compbio.charite.de/hudson/</a> ? website

**Value**

List of HPO terms per MPO term

---

get.node.friendly.long.names  
*Split up the HPO term descriptions so they fit in nodes for plot*

---

**Description**

Split up the HPO term descriptions so they fit in nodes for plot

**Usage**

```
get.node.friendly.long.names(hpo.terms, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms

**Value**

Character vector

**Examples**

```
data(hpo.terms)  
get.node.friendly.long.names(hpo.terms, c("HP:0001873", "HP:0011877"))
```

---

get.ontology	<i>Get R-Object representation of ontology from obo file</i>
--------------	--

---

**Description**

Get R-Object representation of ontology from obo file

**Usage**

```
get.ontology(file, qualifier = "HP")
```

**Arguments**

file	File path of obo file
qualifier	Character vector - "HP" for HPO, "MP" for MPO, etc.

**Value**

R-Object (list) representing ontology

---

get.pop.frequency.based.colours	<i>Function to colour HPO nodes in plot with colours based on information content/frequency of terms with respect to population</i>
---------------------------------	---

---

**Description**

Function to colour HPO nodes in plot with colours based on information content/frequency of terms with respect to population

**Usage**

```
get.pop.frequency.based.colours(hpo.terms, terms, plotting.context,  
  colourPalette = colorRampPalette(c("Yellow", "Green", "#0099FF"))(10),  
  terms.freq = if (is.null(plotting.context$frequency))  
  exp(-plotting.context$information[terms]) else plotting.context$frequency,  
  max.colour.freq = max(terms.freq), min.colour.freq = min(terms.freq))
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms
colourPalette	Character vector of colours for the different information contents of the terms to be plotted, going from rare to common
terms.freq	Numeric vector of frequencies of terms in plot, named by term
max.colour.freq	Numeric value in [0, 1] giving the maximum frequency (to which the dullest color will be assigned)
min.colour.freq	Numeric value in [0, 1] giving the minimum frequency (to which the brightest color will be assigned)

**Value**

Character vector of colours, named by term

---

get.shortened.names     *Get human readable, shortened (where possible) HPO term names*

---

**Description**

Get human readable, shortened (where possible) HPO term names

**Usage**

```
get.shortened.names(hpo.terms, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms

**Value**

Character vector

**Examples**

```
data(hpo.terms)
get.shortened.names(hpo.terms, c("HP:0001873", "HP:0011877"))
```

get.significance.based.sizes

*Function to size HPO nodes in plot with colours based on significance of seeing this many of each term in phenotypes*

---

### **Description**

Function to size HPO nodes in plot with colours based on significance of seeing this many of each term in phenotypes

### **Usage**

```
get.significance.based.sizes(hpo.terms, terms, plotting.context)
```

### **Arguments**

hpo.terms      R-Object representation of HPO  
terms          Character vector of HPO terms  
plotting.context      List object with hpo.phenotypes slot for list of character vectors of terms

### **Value**

Character vector of sizes, named by term

---

get.simple.node.labels

*Function to label HPO nodes in plot with just node description*

---

### **Description**

Function to label HPO nodes in plot with just node description

### **Usage**

```
get.simple.node.labels(hpo.terms, terms, plotting.context)
```

### **Arguments**

hpo.terms      R-Object representation of HPO  
terms          Character vector of HPO terms  
plotting.context      List object with hpo.phenotypes slot for list of character vectors of terms

### **Value**

Character vector of labels, named by term

---

`get.term.adjacency.matrix`*Get an adjacency for set of HPO terms*

---

**Description**

Get an adjacency for set of HPO terms

**Usage**

```
get.term.adjacency.matrix(hpo.terms, terms)
```

**Arguments**

<code>hpo.terms</code>	R-Object representation of HPO
<code>terms</code>	Character vector of HPO terms

**Value**

A logical matrix representing the directed adjacency matrix of terms based on DAG structure of HPO, whereby a TRUE entry signifies that the term corresponding to the column is a parent term of the term corresponding to the row.

**See Also**

[get.term.pseudo.adjacency.matrix](#)

**Examples**

```
data(hpo.terms)
get.term.adjacency.matrix(hpo.terms, c("HP:0000118", "HP:0001873", "HP:0011877"))
```

---

`get.term.descendancy.matrix`*Get logical descendancy matrix for set of terms*

---

**Description**

Get logical descendancy matrix for set of terms

**Usage**

```
get.term.descendancy.matrix(hpo.terms, terms = NULL, rows = terms,
  cols = terms)
```

**Arguments**

<code>hpo.terms</code>	R-Object representation of HPO
<code>terms</code>	Character vector of HPO terms
<code>rows</code>	Rows for resultant matrix (defaults to terms)
<code>cols</code>	Cols for resultant matrix (defaults to terms)

**Value**

A logical descendancy matrix of terms by terms based on DAG structure of HPO, where by the row term is an ancestor of the column term if `result[row.term,col.term] == TRUE`

**Examples**

```
data(hpo.terms)
get.term.descendancy.matrix(hpo.terms, c("HP:0001873", "HP:0011877"))
```

---

`get.term.frequencies` *Get frequency of each term in a set of phenotypes*

---

**Description**

Get frequency of each term in a set of phenotypes

**Usage**

```
get.term.frequencies(hpo.terms, hpo.phenotypes, patch.missing = FALSE)
```

**Arguments**

<code>hpo.terms</code>	R-Object representation of HPO
<code>hpo.phenotypes</code>	List of HPO term character vectors
<code>patch.missing</code>	Logical indicating whether to include all HPO even if they're not present in the <code>hpo.phenotypes</code> as if they had occurred once

**Value**

Numeric vector of information contents, named by corresponding HPO terms. Takes into account ancestors, in the sense that all ancestor terms implied by the phenotypes are considered 'on'

**See Also**

[get.term.info.content](#)

**Examples**

```
data(hpo.terms)
get.term.frequencies(hpo.terms, list("HP:0001873"))
```

---

get.term.info.content *Get information content of each term in a set of phenotypes*

---

**Description**

Get information content of each term in a set of phenotypes

**Usage**

```
get.term.info.content(hpo.terms, hpo.phenotypes, patch.missing = FALSE)
```

**Arguments**

hpo.terms	R-Object representation of HPO
hpo.phenotypes	List of HPO term character vectors
patch.missing	Logical indicating whether to include all HPO even if they're not present in the hpo.phenotypes as if they had occurred once

**Value**

Numeric vector of information contents, named by corresponding HPO terms. Takes into account ancestors, in the sense that all ancestor terms implied by the phenotypes are considered 'on'

**Examples**

```
data(hpo.terms)
get.term.info.content(hpo.terms, list("HP:0001873"))
```

---

get.term.pseudo.adjacency.matrix

*Get an adjacency to MRCA matrix for set of HPO terms*

---

**Description**

Get an adjacency to MRCA matrix for set of HPO terms

**Usage**

```
get.term.pseudo.adjacency.matrix(hpo.terms, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms

**Value**

A logical matrix representing the directed adjacency matrix of terms based on DAG structure of HPO, whereby a TRUE entry signifies the term corresponding to the column is MRCA of the row term in terms

**See Also**

[get.term.adjacency.matrix](#)

**Examples**

```
data(hpo.terms)
get.term.pseudo.adjacency.matrix(hpo.terms, c("HP:0000118", "HP:0001873", "HP:0011877"))
```

---

hpo.plot

*Plot HPO graph object*

---

**Description**

Plot HPO graph object

**Usage**

```
hpo.plot(hpo.terms, terms = apply.term.filters(hpo.terms = hpo.terms,
  plotting.context = plotting.context, term.filters = list()),
  plotting.context = NULL, hpo.phenotypes = NULL, term.frequencies = NULL,
  colours = "cyan", labels = get.simple.node.labels,
  borders = "#FFFFFF00", sizes = 0.75, font.sizes = rep(30,
  length(terms)), shapes = rep("circle", length(terms)), nodeAttrs = NULL,
  arrowAttrs = list(color = "#000000"), ...)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms
hpo.phenotypes	List of HPO term character vectors
term.frequencies	Numeric vector of population frequencies of terms (named by term codes)
colours	Function to set the colours of the HPO nodes in the graph based on the plotting context, or a character vector of colours
labels	Function to set the labels of the HPO nodes in the graph based on the plotting context, or a character vector of node labels
borders	Function to set the borders of the HPO nodes in the graph based on the plotting context, or a character vector of border colours

sizes	Function to set the sizes of the HPO nodes in the graph based on the plotting context, or a numeric vector of node sizes
font.sizes	Function to set the font sizes of the text to be placed in the HPO nodes in the graph based on the plotting context, or an integer vector of font sizes
shapes	Function to set the shapes of the HPO nodes in the graph based on the plotting context, or a character vector of shape names (defaults to 'circle')
nodeAttrs	Pass nodeAttrs directly to rgraphviz plotting function
arrowAttrs	List of properties to set for arrows (note, these properties will be used for all arrow)
...	Extra arguments to pass to plot

**Value**

Plots graph

**See Also**

[get.hpo.graph](#)

**Examples**

```
data(hpo.terms)
hpo.plot(
  hpo.terms=hpo.terms,
  terms=get.ancestors(hpo.terms,
c("HP:0001382", "HP:0004272", "HP:0007917", "HP:0004912", "HP:0001596"))
)
```

---

hpo.terms

*HPO Terms object (based on version 887 of the HPO)*

---

**Description**

Object comprising list of properties of the HPO, indexed by term ID

**Format**

List of indices containing metadata and structure of HPO

---

hpoPlot

*Functions for Plotting HPO Terms*

---

**Description**

Functions for performing operations on sets of HPO terms (character vectors of HPO term IDs) in the context of the HPO structure, and plotting them with a various options

**Details**

Package: hpoPlot  
Type: Package  
Version: 2.3  
Date: 2015-01-7  
License: GPL (>= 2)

The key function is `hpo.plot`, which plots a set of phenotype terms given their HPO IDs and their ontological relations given by the HPO.

### Author(s)

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### References

‘The Human Phenotype Ontology project: linking molecular biology and disease through phenotype data’, Nucl. Acids Res. (1 January 2014) 42 (D1): D966-D974 doi:10.1093/nar/gkt1026 Westbury, S. K. et al. (2015). Human Phenotype Ontology annotation and cluster analysis to unravel genetic defects in 707 cases with unexplained bleeding and platelet disorders. Genome Medicine. 7 (2015)

---

intersection.with.branches

*Intersect set of terms with branches of HPO*

---

### Description

Intersect set of terms with branches of HPO

### Usage

```
intersection.with.branches(hpo.terms, branch.roots, terms)
```

### Arguments

<code>hpo.terms</code>	R-Object representation of HPO
<code>branch.roots</code>	Character vector of roots of branches you wish to intersect with
<code>terms</code>	Character vector of HPO terms

### Value

Character vector of terms

**Examples**

```
data(hpo.terms)
intersection.with.branches(hpo.terms, "HP:0001872", c("HP:0001873", "HP:0011877"))
```

---

mpo.terms	<i>MPO Terms object</i>
-----------	-------------------------

---

**Description**

Object comprising list of properties of the MPO, indexed by term ID

**Format**

List of indices containing metadata and structure of MPO

---

mpo.to.hpo	<i>Object containing data for mapping between MPO and HPO</i>
------------	---

---

**Description**

List containing cross-species ontology (MPO to HPO) information - character vectors of HPO terms indexed by associated MPO term IDs

**Format**

List of HPO terms per MPO term

---

n.most.frequent.terms	<i>Select most frequently annotated terms from a set of phenotypes</i>
-----------------------	--

---

**Description**

Select most frequently annotated terms from a set of phenotypes

**Usage**

```
n.most.frequent.terms(hpo.terms, terms, plotting.context, n)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms
n	Integer

**Value**

Character vector of length at most n

**See Also**

[remove.terms.with.less.than.n.occurrences](#), [remove.uninformative.for.plot](#)

**Examples**

```
data(hpo.terms)
n.most.frequent.terms(hpo.terms, c("HP:0001873"),
list(hpo.phenotypes=list("HP:0001873", "HP:0001902")), n=2)
```

---

*p.values.for.occurrence.of.term.in.group*

*Get p-values for observing at least as many of each term as have been in phenotypes given information content*

---

**Description**

Get p-values for observing at least as many of each term as have been in phenotypes given information content

**Usage**

```
p.values.for.occurrence.of.term.in.group(hpo.terms, hpo.phenotypes, terms.freq)
```

**Arguments**

<code>hpo.terms</code>	R-Object representation of HPO
<code>hpo.phenotypes</code>	List of HPO term character vectors
<code>terms.freq</code>	Numeric vector of population frequencies of terms

**Value**

Numeric vector of log p-values named by corresponding term

---

prune.branch	<i>Prune all terms descending from given term down to that term and ensure no degeneracy</i>
--------------	--

---

**Description**

Prune all terms descending from given term down to that term and ensure no degeneracy

**Usage**

```
prune.branch(hpo.terms, prune.to.point, terms)
```

**Arguments**

hpo.terms	R-Object representation of HPO
prune.to.point	HPO term which can be included, but whose descendants should be excluded
terms	Character vector of HPO terms

**Value**

Character vector of terms

---

remove.links	<i>Remove terms with exactly one parent and child from plot</i>
--------------	---

---

**Description**

Remove terms with exactly one parent and child from plot

**Usage**

```
remove.links(hpo.terms, terms, plotting.context = NULL)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms

**Value**

Character vector

### See Also

[remove.terms.with.less.than.n.occurrences,n.most.frequent.terms](#)

### Examples

```
data(hpo.terms)
remove.links(hpo.terms, c("HP:0001873"), list(hpo.phenotypes=list("HP:0001873", "HP:0001902")))
```

---

remove.non.pa.terms *Remove terms not descending from phenotypic abnormality*

---

### Description

Remove terms not descending from phenotypic abnormality

### Usage

```
remove.non.pa.terms(hpo.terms, terms, plotting.context)
```

### Arguments

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms

### Value

Character vector

### See Also

[remove.terms.with.less.than.n.occurrences,n.most.frequent.terms](#)

---

```
remove.terms.with.less.than.n.occurrences
```

*Remove terms with less than certain number of occurrences*

---

**Description**

Remove terms with less than certain number of occurrences

**Usage**

```
remove.terms.with.less.than.n.occurrences(hpo.terms, terms, plotting.context, n)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
plotting.context	List object with hpo.phenotypes slot for list of character vectors of terms
n	Integer

**Value**

Character vector

**See Also**

[n.most.frequent.terms](#), [remove.uninformative.for.plot](#)

**Examples**

```
data(hpo.terms)
remove.terms.with.less.than.n.occurrences(hpo.terms,
c("HP:0001873"), list(hpo.phenotypes=list("HP:0001873", "HP:0001902")), 2)
```

---

```
remove.uninformative.for.plot
```

*Remove uninformative terms (fitting plotting filter format)*

---

**Description**

Remove uninformative terms (fitting plotting filter format)

**Usage**

```
remove.uninformative.for.plot(hpo.terms, terms = get.ancestors(hpo.terms,
  unlist(plotting.context$hpo.phenotypes)), plotting.context)
```

**Arguments**

hpo.terms      R-Object representation of HPO  
terms          Character vector of HPO terms  
plotting.context  
                 List object with hpo.phenotypes slot for list of character vectors of terms

**Value**

Character vector

**See Also**

[remove.terms.with.less.than.n.occurrences](#), [n.most.frequent.terms](#)

**Examples**

```
data(hpo.terms)
remove.uninformative.for.plot(hpo.terms,
c("HP:0001873"), list(hpo.phenotypes=list("HP:0001873", "HP:0001902")))
```

---

remove.uninformative.terms

*Get a minimal set of terms which can be used to partition a set of phenotypes*

---

**Description**

Get a minimal set of terms which can be used to partition a set of phenotypes

**Usage**

```
remove.uninformative.terms(hpo.terms, hpo.phenotypes)
```

**Arguments**

hpo.terms      R-Object representation of HPO  
hpo.phenotypes List of HPO term character vectors

**Value**

Character vector of set of terms, excluding terms for which the presence of their descendants all partition the set of terms in the same way

**Examples**

```
data(hpo.terms)
remove.uninformative.terms(hpo.terms, list(Patient1=c("HP:0001873")))
```

---

setDimNames	<i>setNames for arrays...</i>
-------------	-------------------------------

---

**Description**

setNames for arrays...

**Usage**

```
setDimNames(array.object, list.of.dimension.names)
```

**Arguments**

array.object    Array

list.of.dimension.names

List of character vectors with which to name each dimension of the array

**Value**

Named array

**Examples**

```
setDimNames(matrix(1:4,2,2), list(c("Cat", "Dog"), c("Name", "Weight")))
```

---

simpleCap	<i>Capitalise words in character vector</i>
-----------	---

---

**Description**

Capitalise words in character vector

**Usage**

```
simpleCap(x)
```

**Arguments**

x                    Character vector

**Value**

Character vector

**Examples**

```
simpleCap(c("a simple test", "Another-test"))
```

---

swap.out.alt.ids	<i>Remove alternate/deprecated HPO term IDs and swap for new ones</i>
------------------	---

---

**Description**

Remove alternate/deprecated HPO term IDs and swap for new ones

**Usage**

```
swap.out.alt.ids(hpo.terms, terms, remove.dead = FALSE)
```

**Arguments**

hpo.terms	R-Object representation of HPO
terms	Character vector of HPO terms
remove.dead	Boolean to indicate whether to strip out terms which can't be found in the given hpo.terms database argument

**Value**

A directed adjacency matrix of terms based on DAG structure of HPO, whereby each term is considered adjacent to it's MRCA in terms

**Examples**

```
data(hpo.terms)
swap.out.alt.ids(hpo.terms, c("HP:0001873"))
```

---

```
term.set.list.from.character
```

*Get list of character vector of HPO terms, given character vector of comma separated terms*

---

**Description**

Get list of character vector of HPO terms, given character vector of comma separated terms

**Usage**

```
term.set.list.from.character(character.vector)
```

**Arguments**

character.vector	Character vector of comma separated terms
------------------	---

**Value**

List of character vectors of HPO terms

**Examples**

```
term.set.list.from.character(c("HP:0001873", "HP:0001902"))
```

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